

B160

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
9 January 2003 (09.01.2003)

PCT

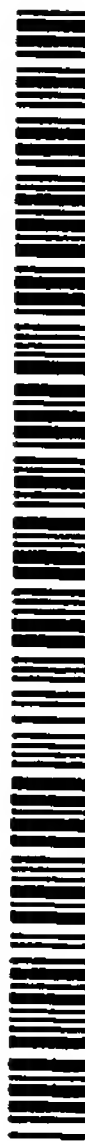
(10) International Publication Number
WO 03/002767 A1

- (51) International Patent Classification⁷: C12Q 1/68, (74) Agent: JOYCE, Timothy, H.; Agilent Technologies, Inc.,
C12P 19/34, C07H 21/04 Intellectual Property Administration, M/S DL429, P.O.
Box 7599, Loveland, CO 80537-0599 (US).
- (21) International Application Number: PCT/US02/14193 (81) Designated State (*national*): JP.
- (22) International Filing Date: 2 May 2002 (02.05.2002) (84) Designated States (*regional*): European patent (AT, BE,
CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC,
NL, PT, SE, TR).
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data:
09/896,299 29 June 2001 (29.06.2001) US
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Published:

— with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



WO 03/002767 A1

(54) Title: METHOD OF DNA SEQUENCING USING CLEAVABLE TAGS

(57) Abstract: The present invention provides novel systems for sequencing nucleic acid molecules using dNTPs that are 3' end labeled with cleavable tags that block further extension and uniquely identify the bases to which they are attached. Removal of the tags liberates the 3' ends of the extension products for further extension. In related embodiments, oligonucleotides containing sequence-related cleavable tags are employed in a ligation reaction to determine the sequence of a particular DNA sample.

1 METHOD OF DNA SEQUENCING USING CLEAVABLE TAGS

2 Background of the Invention

3 DNA sequencing is an important analytical technique critical to generating
4 genetic information from biological organisms. The increasing availability of rapid
5 and accurate DNA sequencing methods has made possible the determination of the
6 DNA sequences of entire genomes, including the human genome. DNA sequencing
7 has revolutionized the field of molecular biological research. In addition, DNA
8 sequencing has become an important diagnostic tool in the clinic, where the rapid
9 detection of a single DNA base change or a few base changes can be used to detect,
10 for example, a genetic disease or cancer.

11 Most current methods of DNA sequencing are based on the method of Sanger
12 (*Proc. Natl. Acad. Sci. U.S.A.*, 74, 5463 (1977)). This method relies on gel
13 electrophoresis of single stranded nucleic acid fragments that are generated when a
14 polymerization extension reaction of a primer is terminated by incorporation of a
15 radioactively labeled dideoxynucleotide triphosphate. Short strands of DNA are
16 synthesized under conditions that produce DNA fragments of variable length using a
17 DNA polymerase and deoxynucleotide triphosphates (dNTP). A small amount of
18 dideoxynucleotide triphosphates (ddNTP) is introduced into the DNA synthesis
19 mixture so that chain terminating ddNTPs are sometimes integrated into a growing
20 strand. Typically, four different extension reactions are performed side by side, each
21 including a small amount of one ddNTP. Each extension reaction produces a mixture
22 of DNA fragments of different lengths terminated by a known ddNTP. The ratio of
23 ddNTPs to dNTPs is chosen so that the populations of DNA fragments in any given

1 extension reaction includes fragments of all possible lengths (up to some maximum)
2 terminating with the relevant ddNTP. The nucleic acid fragments are separated by
3 length in the gel, typically utilizing a different lane in a polyacrylamide gel for each of
4 the four terminating nucleotide bases being detected. However, such size exclusion
5 chromatography is generally a low resolution method limited to reading short
6 sequences.

7 A variation of this method utilizes dyes rather than radioactivity to label the
8 ddNTPs. Different dyes are used to uniquely label each of the different ddNTPs (i.e.,
9 a different dye may be associated with each of A, G, C, and T termination) (Smith et
10 al. and Prober et al. *Science* 238:336-341, 1987). In the method of Smith, fluorescent
11 dyes are attached to the 3' end of the dNTP converting it into a ddNTP. The use of
12 four different dye labels allows the entire sequencing reaction to be conducted in a
13 single reaction vessel and results in a more uniform signal response for the different
14 DNA fragments. The dye-terminated dNTPs are also able to be electrophoresed in a
15 single lane. The advent of capillary electrophoresis further increased the separation
16 efficiency of this method, allowing shorter run times, longer reads, and higher
17 sensitivity.

18 Despite these advances, DNA sequencing methods that rely on electrophoresis
19 to resolve DNA fragments according to their size are limited by the rate of the
20 electrophoresis and the number of bases that are detectable on the gel. In addition,
21 real time imaging of the gel is not possible. Accordingly, in order to increase the
22 speed and reliability of the sequencing reaction, great effort has been made to
23 automate these steps. Automated DNA sequencing machines are now available that
24 are capable of high throughput sequencing for both genomic sequencing and routine

1 clinical applications. However, these newer techniques remain cumbersome,
2 requiring specialized chemicals and the intensive labor of skilled technicians.

3 One newer method of DNA sequencing, "pyrosequencing" or "sequencing-by-
4 synthesis," disclosed in WO 98/13523, is based on the concept of detecting inorganic
5 pyrophosphate (PPi), which is released during a polymerase reaction. As in the
6 Sanger method, a sequencing primer is hybridized to a single stranded DNA template
7 and incubated with a DNA polymerase. In addition to the polymerase, the enzymes
8 ATP sulfurylase, luciferase, and apyrase, and the substrates, adenine 5'
9 phosphosulfate (APS) and luciferin, are added to the reaction. Subsequently,
10 individual nucleotides are added. When the added nucleotide is complementary to the
11 next available base in the template strand, it is incorporated into the extension
12 product. Such incorporation of a complementary base is accompanied by release of
13 pyrophosphate (PPi), which is converted to ATP in the presence of adenosine 5'
14 phosphosulfate by apyrase in a quantity equimolar to the amount of incorporated
15 nucleotide. The ATP generated by the reaction with apyrase then drives the luciferase
16 mediated conversion of luciferin to oxyluciferin, generating visible light in amounts
17 that are proportional to the amount of ATP and thus the number of nucleotides
18 incorporated into the growing DNA template. The light produced by the luciferase-
19 catalyzed reaction is detected by a charge coupled device (CCD) camera and detected
20 as a peak in a pyrogram™.

21 In a pyrosequencing reaction, if the first nucleotide added to the reaction is not
22 complementary to the next available nucleotide on the growing DNA strand there is
23 no light generated. If no light is generated by the addition of the first nucleotide, a
24 second of four dNTPs is added sequentially to the reaction to test whether it is the

1 complementary nucleotide. This process is continued until a complementary
2 nucleotide is added and detected by a positive light read-out. Whether or not a
3 positive light reaction is generated, apyrase, a nucleotide-degrading enzyme,
4 continuously degrades unincorporated dNTPs and excess ATP in the reaction mixture.
5 When degradation is complete, another dNTP is added.

6 Although pyrosequencing is capable of generating high quality data in a
7 relatively simple fashion, this method has several drawbacks. First, the productivity
8 of the method is not high, reading only about 1 base per 100 seconds. The rate of the
9 reaction is limited by the necessity of having to add new enzymes with each addition
10 of the dNTPs in addition to the necessity of having to test each of the four dNTPs
11 separately. In addition, it has been found that the dATP used in the chain extension
12 reaction interferes in subsequent luciferase-based detection reactions by acting as a
13 substrate for the luciferase enzyme. Finally, these reactions are expensive to run.

14 While pyrosequencing improves the ease and speed with which DNA
15 sequencing is achieved, there exists the need for improved sequencing methods that
16 allow more rapid detection. Preferred techniques would be amenable to automation
17 and allow the sequence information to be revealed simultaneously with or shortly
18 after the chain extension reaction.

19 **Summary of the Invention**

20 The present invention provides a novel system for sequencing nucleic acid
21 molecules. In particular, the invention utilizes dNTPs that are 3' end labeled with a
22 cleavable tag that distinguishes the dNTP from other dNTPs (e.g., the tag may be
23 unique to the dNTP). The cleavable tags are functional groups that can be later

1 removed by any appropriate means, including but not limited to, exposure to chemical
2 cleavage conditions or light. dNTPs labeled with the cleavable tags function as
3 terminated dNTPs (cdNTPs), in that their incorporation into a single stranded nucleic
4 acid molecule via a primer extension reaction blocks further extension. However,
5 removal of the tag converts the cdNTP back into an extendible nucleotide.

6 According to the present methods, a sequencing primer is hybridized to a
7 nucleic acid template, e.g., a single stranded DNA template, and incubated with an
8 enzyme (DNA polymerase) and four cdNTPs (tag terminated dATP (cdATP), dCTP
9 (cdCTP), dGTP (cdGTP), and dTTP (cdTTP)). The DNA polymerase then extends
10 the primer by adding to it whichever cdNTP is complementary to the next available
11 base on the template strand. Only a single cdNTP is incorporated, because the cdNTP
12 cannot be further extended.

13 After completion of a single base addition, unreacted (excess) cdNTPs are
14 removed from the reaction mixture, which includes the extended primer, the DNA
15 polymerase, and the single stranded DNA template. The step of removing can be
16 accomplished by any of a variety of means that would be apparent to one skilled in
17 the art. For example, if the reaction mixture is contained in a chamber that has an
18 attached membrane (e.g., an ultrafiltration membrane that allows small molecules
19 such as water, salts, and cdNTPs to pass through, but does not allow passage of large
20 molecules such as single stranded DNA), the excess cdNTP can be washed through
21 the membrane. Alternatively, if the single stranded DNA is attached to a solid
22 support, the excess cdNTPs can be washed away from the single stranded DNA
23 without dislodging the hybridized, extended primer.

1 Once the step of removing is complete, the tag is cleaved from the cdNTP that
2 is extended into the single stranded DNA template. In certain embodiments, the
3 cleavage occurs by photo-cleavage of the tag from the extended single stranded DNA
4 template by exposure to light. Alternatively, in other preferred embodiments, the
5 cleavage occurs by exposure of the single stranded DNA template to a chemical
6 cleaving agent, e.g., an acid or a base. Whichever cleavage method is employed, the
7 result is liberation of the 3' end of the extension product for further extension.

8 The cleaved tag is then washed through the membrane into a detector for
9 identification, thereby identifying the complementary base in the single stranded
10 DNA template and determining the DNA sequence. The detector used to identify the
11 tag is chosen based on the type of cleavable tag employed. Any of a variety of tags
12 may be employed in the present invention, as would be recognized by the skilled
13 artisan, and such tags are described herein. Once the tag is cleaved, the four cdNTPs
14 are added back to the primer extension reaction mixture and the cycle of extension,
15 tag cleavage, and identification is repeated.

16 In other preferred embodiments, short oligonucleotides are employed in a
17 ligation reaction to determine the sequence of a particular DNA sample. The
18 sequence of a DNA sample is determined by incorporating "X" complementary bases
19 (e.g., 2mers, 3mers, or more) at a time onto the single stranded DNA template
20 adjacent to a primer using a DNA ligase instead of using a DNA polymerase. Each
21 oligonucleotide is tagged and labeled with a cleavable tag so that the position of each
22 base in the sequence of the oligonucleotide can be identified. The tag further prevents
23 ligation of the oligonucleotides to one another.

1 According to this aspect of the invention, a template DNA is exposed to the
2 oligonucleotides, the oligonucleotides are allowed to hybridize to the template DNA,
3 and a ligation reaction is allowed to take place on the DNA template such that one
4 complementary oligonucleotide is incorporated onto the DNA template adjacent to the
5 annealed primer. Following ligation, the unincorporated oligonucleotides are washed
6 away from the DNA sample and the tags are cleaved and analyzed to determine the
7 nucleic acid sequence.

8

9 Detailed Description of Certain Preferred Embodiments

10 The present invention provides a system for sequencing a DNA molecule
11 using deoxynucleotide triphosphates of adenine, thymine, guanine, and cytosine that
12 are each labeled with a different cleavable tag that is used to identify the base. In
13 preferred embodiments, the cleavable tag further acts as a terminator to extension of a
14 single stranded DNA template in a polymerase extension reaction until the tag is
15 removed from the incorporated base. Once removed, the tag is isolated and identified,
16 and the process of base addition and cleavage is repeated. More particularly, the steps
17 of extension, cleavage, and detection are repeated until sufficient sequence of the
18 single stranded DNA template is determined.

19 According to certain preferred embodiments, inventive methods of
20 determining the sequence of a nucleic acid include the steps of (a) hybridizing an
21 oligonucleotide to a single stranded DNA, wherein the oligonucleotide is
22 complementary to at least a portion of the single stranded DNA; (b) providing a DNA
23 polymerase and four deoxynucleotide triphosphates (dNTPs) (e.g., dATP, dGTP,

1 dCTP, and dTTP) wherein each dNTP is 3' end labeled with a cleavable tag (cdNTP)
2 that distinguishes the dNTP from other dNTPs; (c) extending the single stranded DNA
3 hybridized to the oligonucleotide by adding one complementary cdNTP in a
4 polymerase extension reaction, wherein the tag on the extended cdNTP blocks further
5 extension by the DNA polymerase; (d) optionally removing excess cdNTPs that are
6 not extended onto the single stranded DNA; (e) cleaving the tag from the extended
7 cdNTP; and (f) detecting the tag so that the incorporated base is detected. In certain
8 preferred embodiments, the method includes the step of repeating steps (a) through (f)
9 on the sample of single stranded DNA.

10 As indicated above, prior to cleavage of the tag from the extended base on the
11 DNA template, the excess, unincorporated cdNTPs are preferably removed from the
12 extension reaction. According to the invention, the tags may be removed by any of a
13 variety of washing or rinsing procedures that separate the excess, unincorporated
14 dNTPs from the extended DNA template. In one preferred embodiment, the
15 extension reaction is contained within a chamber that has an attached filtration
16 membrane, e.g., an ultrafiltration membrane, that allows small molecules such as
17 water, salts, and cdNTPs to pass through, while retaining large molecules such as
18 ssDNA. According to this particular embodiment, a wash solution, e.g., a buffered
19 saline solution such as phosphate buffered saline, is passed through the ultrafiltration
20 membrane of the chamber containing the oligonucleotide primer, the DNA
21 polymerase, the cdNTPs, and the extended DNA to rinse away the excess cdNTPs.
22 Alternatively, if the DNA template is attached to a solid support, a wash solution may
23 be passed over the solid support to rinse the excess cdNTPs away from the solid
24 support.

1 In a related embodiment, the sequencing method of the present invention is
2 also amenable to sequence determination via oligonucleotide ligation. This technique
3 requires first exposing the DNA template to a collection of tagged oligonucleotides
4 (e.g., the tagged oligonucleotides may be a collection of short randomized
5 oligonucleotides). Preferably, a 3' tag blocks further ligation at the 3' end of the
6 oligonucleotide to other oligonucleotides in the collection. However, it will be
7 appreciated that if the tag is located at a position on the oligonucleotide other than the
8 3' end, the 3' end of the oligonucleotide would still need to be blocked, for example,
9 with another functional group. Once the DNA template is mixed with the tagged
10 oligonucleotides, the oligonucleotides are allowed to hybridize to the DNA template
11 in a position adjacent to an oligonucleotide primer so that the oligonucleotide and
12 primer can be ligated. Unligated oligonucleotides are then rinsed away from the DNA
13 template, tags are cleaved from the ligated oligonucleotides, and cleaved tags
14 representing the bases of the ligated oligonucleotide are detected. This cycle can be
15 repeated as described, with addition of the oligonucleotide mix occurring at each
16 repetition.

17 In certain preferred embodiments, the number of tags attached to the 3' end of
18 the oligonucleotide may be based on the sequence length of the oligonucleotide. For
19 example, an oligonucleotide that is three bases long may be 3' end labeled with three
20 tags that are attached in a sequential order matching the sequential order of the bases
21 of the oligonucleotide.

22 In the oligonucleotide ligation reaction, as with the polymerase reaction, the
23 DNA template is a single stranded DNA template that is annealed to a primer for
24 primer extension. By "single stranded DNA template" is meant any single stranded

1 DNA template or single stranded DNA template that is partially single stranded, i.e.,
2 may be partially double stranded. In one preferred embodiment, an oligonucleotide
3 that is 3' end blocked and complementary to the sequence adjacent to the primer
4 anneals to the DNA template and is joined to the adjacent primer via a ligase (e.g., T4
5 DNA ligase). The tags on the complementary oligonucleotide are then removed for
6 detection and identification, freeing the 3' end of the complementary oligonucleotide
7 for subsequent rounds of ligation. In such subsequent rounds, the ligase joins the next
8 complementary blocked oligonucleotide to the 3' end of the previously extended
9 primer and the cycle repeats.

10 As mentioned above, the collection of oligonucleotides may include short
11 randomized oligonucleotides. Those skilled in the art will appreciate that the longer
12 the oligonucleotide, the greater the number of oligonucleotides will have to be
13 generated to encompass all possible random oligonucleotide sequences, based on
14 randomization between four bases at each position of the oligonucleotide. For
15 example, generation of a collection of 2mers that encompasses all possible 2mers
16 would require sixteen oligonucleotide sequences; generation of a collection of 3mers
17 that encompasses all possible 3mers would require a panel of 64 oligonucleotide
18 sequences; 4mers would require a panel of 256 oligonucleotide sequences, etc.

19 Identification of an optimal oligonucleotide length may require simply testing various
20 short random oligonucleotide mixes and determining which give the most rapid and
21 accurate DNA sequencing results via oligonucleotide ligation. Of course, the longer
22 the oligonucleotide, the faster the sequencing reaction will proceed, due to the
23 increased number of incorporated bases detected simultaneously. Using this

1 approach, at each round of the sequencing reaction, the oligonucleotide sequence that
2 is ligated to the primer is detected and identified.

3 In certain preferred embodiments, it is conceivable that only a small subset of
4 all possible oligonucleotides need to be used in the sequencing reaction, for example,
5 if the sequence of the DNA template were partially determined (i.e., if certain
6 positions of the oligonucleotide were fixed, fewer base positions would need to be
7 randomized, limiting the number of oligonucleotide required to include all possible
8 permutations). In this particular embodiment, the oligonucleotides could be longer
9 (e.g., 5mers, 6mers, 7mers, 8mers, 9mers, 10mers, or greater than 15mers, 20mers,
10 25mers, 30mers and higher). It is also possible that in certain circumstances one
11 would not need to use as many tags i.e., one would not need to use one tag for every
12 base. For example, one unique tag could be used to identify an entire oligonucleotide
13 sequence.

14 In other preferred embodiments, two or more unique tags could be used to
15 identify an entire oligonucleotide sequence, the total number of tags being less than
16 the total number of bases in the oligonucleotides (e.g., each tag could identify short
17 sequential stretches of oligonucleotides (e.g., a 3mer or a 4mer etc.) within the entire
18 oligonucleotide sequence). In a related embodiment, an oligonucleotide, particularly
19 an oligonucleotide used in the ligation aspect of the invention, may not be randomized
20 at every position (e.g., if certain nucleotide positions are fixed), and may even be
21 randomized at only one or several positions, e.g., 1-2, 1-3, 1-4 or 1-5 positions.
22 Under these circumstances, only a subset of possible variations would be relevant.

23 In embodiments where the length of the oligonucleotide sequence increases
24 the number of tags required to identify the oligonucleotide sequence, the availability

1 of many unique mass tags makes mass spectrometry a particularly useful system for
2 detection. Since each short random oligonucleotide must be labeled with a unique
3 tags, the short random oligonucleotide may have a maximum length in certain
4 circumstance (e.g., the length and number of oligonucleotides in a collection of
5 oligonucleotides may be limited by the availability of different unique tags).
6 However, mass tags may have the same nominal mass and vary in structure, thereby
7 increasing the diversity of tags available.

8 Although the level of diversity available in the mass spectrometry system is
9 sufficient to permit unique MS/MS fragmentation, those skilled in the art will
10 appreciate that, because identification of the incorporated oligonucleotides is based on
11 the MS/MS parent/daughter transition, if an MS/MS approach is used, multiplexing
12 target DNA samples is not possible. The MS/MS approach requires the isolation of a
13 single mass followed by fragmentation and mass analysis. Multiplexing would
14 present too many masses for isolation and fragmentation to be practical. However,
15 the MS/MS approach would be helpful in increasing the potential number of mass
16 tags required for coding the oligonucleotides used in the ligation reaction.

17 Thus, according to the oligonucleotide ligation aspect of the invention, the
18 sequence of a single stranded DNA template may be determined by (a) hybridizing a
19 complementary oligonucleotide to a single stranded DNA adjacent to a primer,
20 wherein the oligonucleotide is 3' end labeled with one or more cleavable tags unique
21 to the oligonucleotide sequence; (b) ligating the hybridized complementary
22 oligonucleotide to the primer, wherein the one or more tags on the extended cdNTP
23 blocks further ligation by the DNA ligase; (c) optionally removing excess
24 oligonucleotides that are not ligated; (d) cleaving the one or more tags from the

1 ligated complementary oligonucleotide; (e) detecting the one or more tags. In certain
2 preferred embodiments, steps (a) through (e) are repeated on the single stranded
3 DNA.

4 As in the polymerase reaction, in the ligation reaction, prior to cleavage of the
5 tag(s) from the extended oligonucleotide on the DNA template, the excess
6 oligonucleotides are preferably removed. The oligonucleotides may be removed by
7 any of a variety of separation procedures that may include washing or rinsing the
8 unincorporated oligonucleotides away from the extended DNA template. As with the
9 polymerase reaction, in one preferred embodiment, the ligation reaction is contained
10 within a chamber that has an attached filtration membrane that would allow short
11 oligonucleotides to pass through, while retaining larger molecules, such as the DNA
12 template. Alternatively, the DNA template is attached to a solid support and a wash
13 solution may be passed over the solid support to remove the unincorporated tagged
14 oligonucleotides.

15 As will be appreciated by those skilled in the art, whether the sequencing
16 reaction employs a DNA polymerase or a DNA ligase, any tag that is cleavable by
17 chemical means or by light can be used in the present invention. In certain preferred
18 embodiments, the tag is cleaved by exposure to an acid or a base. In other preferred
19 embodiments, the tag is cleaved by exposure to light, i.e., in a photo-cleavage
20 reaction. The cleavable tags themselves include any functional group that imparts a
21 unique identity onto the oligonucleotide or base that is tagged. According to the
22 present invention, useful tags include, e.g., fluorescent tags, mass tags, IR tags, UV
23 tags, potentiometric tags, etc. For example, a fluorescent tag may be attached to a
24 dNTP prior to the primer extension reaction, and then may be cleaved from the dNTP

1 after the dNTP is incorporated into the extended DNA strand by exposure of the
2 extended DNA strand to an acid, a base, or light, and analyzed using fluorescence
3 spectrometry. As but another example, a base having an acid, base, or light cleavable
4 mass tag, after incorporation into the DNA template, may be cleaved from the
5 extended DNA strand using the appropriate cleavable agent, and then may be
6 analyzed using mass spectrometry.

7 The DNA sequencing methods of the present invention provide an advantage
8 over existing Sanger-based methods by eliminating the need to separate cDNA
9 fragments on a gel, resulting in longer sequence reads. The present method is rapid
10 and fully automatable. In addition, the selection and detection of one of the four
11 bases is carried out simultaneously.

12 Alternatively, the identification step need not be carried out simultaneously
13 with the cycling of the reaction. For example, the tags from each cycle may be
14 collected and pooled (e.g., onto a 96 well plate). Alternatively, the tags from each
15 cycle may be spatially arrayed (e.g., onto a chip) and the positional information used
16 for identification. Using either method, the tags are analyzed subsequent to the
17 cycling reaction by art available means. Such collection and analysis may increase
18 the speed of the sequencing reaction to increase the throughput of the technique. Of
19 course one skilled in the art would recognize that the appropriate instrumentation is
20 required to analyze the collected tags.

21 Certain aspects of the present invention are described in further detail below.

22

23

24

1 *Nucleic Acid Preparation*

2 In certain preferred embodiments of the invention, the DNA sample is a single
3 stranded DNA template. Alternatively, if in a polymerase extension reaction a
4 thermostable DNA polymerase enzyme is employed, the DNA sample may be double
5 stranded.

6 The DNA sample of the invention may be provided from any available source
7 of DNA, including, for example, a biological sample, including not only samples
8 obtained from living organisms (e.g., mammals, fish, bacteria, parasites, viruses,
9 fungi, and the like) or from the environment (e.g., air water, or solid samples), but
10 biological materials which may be artificially or synthetically produced (e.g., phage
11 libraries, organic molecule libraries, pools of genomic clones, and the like).

12 Representative examples of biological samples include biological fluids (e.g., blood,
13 semen, cerebral spinal fluid, urine), biological cells (e.g., stem cells, B or T cells,
14 fibroblasts, and the like), and biological tissues. Alternatively, the DNA may be a
15 cDNA synthesized from an RNA sample (e.g., from a natural or synthetic source).
16 Such cDNA synthesis may be carried out using reverse transcription, and such
17 systems are readily available.

18 The DNA sample, whether from a biological or synthetic source, may further
19 be amplified, particularly if the amount of sample DNA is small. Amplification can
20 be carried out by any art available method, for example, *in vitro* by PCR or Self
21 Sustained Sequence Replication (3SR) or *in vivo* using a vector. Alternatively, if
22 desired, *in vitro* and *in vivo* amplification may be used in combination (see, e.g.,
23 McPherson, "PCR: A Practical Approach," Oxford University Press, New York,

1 1991). Within other embodiments of the invention, the DNA samples of the present
2 invention may be generated by, for example, a ligation or cleavage reaction.

3 According to the invention, the DNA sample, amplified or unamplified, is
4 either immobilized on a solid support or in solution. In the case of an amplified DNA
5 sample, those skilled in the art will recognize that any amplification procedure may be
6 modified to allow for attachment of the amplified DNA sample to a solid support. For
7 example, a chosen PCR primer may be immobilized to a solid support or may be
8 provided with a means for attachment to a solid support. Immobilization may take
9 place as part of a PCR amplification, e.g., where one or more primers is attached to a
10 support. Alternatively, one or more primers may carry a functional group, e.g., a
11 biotin or thiol group, permitting subsequent immobilization of the DNA sample.
12 Immobilization of the 5' end of a DNA in the sample, e.g., via a 5' primer, allows the
13 DNA to be attached to a solid support, leaving its 3' end remote from the support and
14 available for subsequent hybridization with the extension primer and extension by the
15 polymerase (or ligase). Alternatively, an unamplified DNA sample, such as a vector
16 or a biological sample, may include, or be modified to include, a functional group that
17 allows attachment to a solid support. In a related embodiment, the vector may include
18 a means for attachment to a solid support adjacent to the site of insertion of the
19 sample DNA such that the amplified DNA sample and the means for attachment may
20 be excised together.

21 The solid support may conveniently take the form of, for example, microtiter
22 wells, a solid support activated with polystyrene to bind the DNA sample (e.g., primer
23 DNA), particles, beads (e.g., nylon beads, polystyrene microbeads, or glass beads)
24 (Polysciences, Warrington, PA), glass surfaces, plates, dishes, flasks (Corning Glass

1 Works, Corning, NY), meshes (Becton Dickinson, Mountain View CA), membranes
2 (Millipore Corp., Bedford, MA), dipsticks, capillaries, hollow fibers (Amicon
3 Corporation, Danvers, MA), screens and solid fibers (Edelman et al., U.S. Patent No.
4 3,843,324; see also Kuroda et. al., U.S. Patent No. 4,416,777, incorporated herein by
5 reference), or needles, made, for example, of agarose, cellulose, alginate, Teflon, or
6 polystyrene. Magnetic particles, e.g., majestic beads, may also be used as solid
7 supports, and such materials are commercially available (Robbin Scientific, Mountain
8 View, CA).

9 The solid support may alternatively or additionally carry functional groups
10 such as hydroxyl, carboxyl, aldehyde, or amino groups, or other moieties, such as
11 avidin or streptavidin, for the attachment of the appropriately modified DNA, e.g., via
12 modified oligonucleotide primers used in an amplification reaction. These may in
13 general be provided by treating the support to provide a surface coating of a polymer
14 carrying one of such functional groups, e.g., polyurethane together with a polyglycol
15 to provide hydroxyl groups, or a cellulose derivative to provide hydroxyl groups, a
16 polymer or copolymer of acrylic acid or methacrylic acid to provide carboxyl groups,
17 or an aminoalkylated polymer to provide amino groups. Various other supports and
18 methods of attachment and detachment of nucleic acid molecules to supports, with
19 and without the use of a linker, is described in U.S. Patent No. 5,789,172,
20 incorporated herein by reference.

21 As indicated above, the DNA sample need not be attached to a solid support.
22 For example, a polymerase extension reaction may be carried out in solution on a
23 DNA sample that is prepared in the context of a primer extension reaction having a
24 buffer that will accommodate the addition of an oligonucleotide primer, a DNA

1 polymerase, cdNTPs, and a single or double-stranded DNA template. A ligation
2 extension reaction may be similarly carried out in an appropriate buffer in the
3 presence of an oligonucleotide primer, a DNA ligase, tagged oligonucleotide, and a
4 single or double-stranded DNA template.

5

6 *Extension*

7 Once a suitable DNA sample is prepared, the sample is subject to a primer
8 extension reaction by addition of an oligonucleotide primer, a DNA polymerase, and
9 four cdNTPs, such that one base is incorporated onto the DNA template before
10 extension is blocked by the cleavable tag on the incorporated base. Alternatively, an
11 oligonucleotide ligation reaction is used to extend the template DNA sample, as
12 described above. Those skilled in the art will appreciate that such extension reactions
13 can be modified to accommodate variations in template DNAs, reaction conditions,
14 etc. It will be further recognized that the chosen oligonucleotide primer must be
15 sufficiently large to provide appropriate hybridization with the target DNA sequence.
16 Moreover, the oligonucleotide primer preferably hybridizes immediately 5' to the
17 target sequence. Guidance for selection of primers and primer extension reactions can
18 be found in the scientific literature, for example, Maniatis et al., *Molecular Cloning, a*
19 *laboratory Manual* (1989).

20 The polymerase in the primer extension reaction may be any polymerase that
21 incorporates dNTPs, and preferably cdNTPs, onto a single stranded DNA template.
22 Examples of suitable polymerases that may conveniently be used, and many are
23 known in the art and reported in the literature, include T7 polymerase, Klenow, and
24 Sequenase. Those skilled in the art will be aware that certain polymerases, e.g., T7

1 polymerase, recognize a specific leader sequence in the DNA, which can be included
2 in the sequence of the oligonucleotide primer. If a double stranded DNA template is
3 to be used in the polymerase extension reaction, it is desirable that a thermostable
4 polymerase, such as a Taq polymerase, be chosen to permit repeated temperature
5 cycling without having to add additional polymerase for each round of extension.

6 It is well known that many polymerases have a proof-reading or error
7 checking ability, which sometimes results in digestion of 3' ends available for
8 extension. In the method of the invention, such digestion may result in an increased
9 level of background noise. In order to avoid this problem, a nonproof-reading
10 polymerase, e.g., an exonuclease deficient (exo-) Klenow polymerase may be used.
11 Otherwise, fluoride ions or nucleotide monophosphates that suppress 3' digestion by
12 the polymerase may be added to the extension reaction mixture. In addition, it may
13 be advantageous to use an excess amount of polymerase over primer/template to
14 maximize the number of free 3' ends that are extended. Those skilled in the art will
15 appreciate that the precise reaction conditions and concentrations of reactants etc. may
16 readily be determined for each system according to choice.

17 Since the primer is extended by a single base (or a single oligonucleotide) by
18 the methods described above, the extended primer serves in exactly the same way in
19 the repeated procedure, and with each subsequent base (or oligonucleotide) addition,
20 to determine the next base or bases in the sequence, permitting the whole sample to be
21 sequenced.

22

1 *Separation*

2 In the case of the polymerase extension reaction, prior to cleavage of the tag
3 from the extended DNA template, the excess cdNTPs must be removed from the
4 reaction mixture to prevent contamination of the cleavage product with signals from
5 other unincorporated bases. As mentioned above, this separation may be
6 accomplished by washing the cdNTPs through a membrane filter that allows flow
7 through of small molecules such as water, salts, and cdNTPS, but does not allow the
8 flow through of larger molecules such as the polymerase and the DNA template.

9 In the case of the ligase extension reaction, prior to cleavage of the tag from
10 the oligonucleotide on the extended DNA template, the unincorporated tagged
11 oligonucleotide must be removed from the reaction mixture to prevent contamination
12 of the cleavage product with signals from the unincorporated tagged oligonucleotides.

13 Depending on whether the DNA template is free in solution or attached to a solid
14 support, the excess unincorporated tagged oligonucleotide may be removed by either
15 filtration or washing the solid support, respectively. Those skilled in the art will
16 appreciate that if the DNA ligase is removed from the extension reaction mixture
17 along with the tagged oligonucleotide, the ligase will need to be added back to the
18 extension reaction mixture in subsequent rounds. This, of course, is also applicable to
19 a sequencing reaction that utilizes a polymerase, where the polymerase is removed
20 from the extension reaction in a separate step with the cdNTPs.

21 Those skilled in the art will further appreciate that a wide variety of membrane
22 filters are available in the art. For example, molecular filtration, also known as
23 ultrafiltration, is a membrane separation technique used to segregate substances
24 according to molecular weight and size. Molecular filtration is ideally suited to

1 separate salts and other low molecular weight solutes from high molecular weight
2 species. Molecular filtration is based on a pressure differential across the
3 semipermeable membrane to drive permeable materials through the membrane. For
4 this reason, molecular filtration typically separates solutes and concentrates retained
5 materials more rapidly. Molecular filtration membranes appropriate for use in the
6 present invention may be purchased from Millipore Corp., Bedford, MA.

7 In another preferred embodiment, a flow through cell is used for single
8 stranded DNA analysis. In this embodiment, the tag is washed away and is sent to the
9 detector directly. One example of a variation on the flow through cell approach that
10 would be amenable to multiplexing is to use a 96 well plate with an ultrafiltration
11 membrane incorporated in the well. The excess reagents are either washed through by
12 pressure or centrifuged through. The tag is then subsequently cleaved from extended
13 nucleotide base, washed through the membrane, and collected for analysis by the
14 method appropriate for the type of tag to be identified. In certain preferred
15 embodiments, the different wells are pooled and the tags analyzed simultaneously to
16 provide greater sample multiplexing as well as throughput.

17 Where the DNA template is immobilized on a solid support, the separation is
18 accomplished by simply washing the cdNTPs (or tagged oligonucleotide) away from
19 the solid support. For example, one basic approach to retaining the DNA for analysis
20 would be to absorb the target DNA to an adsorptive surface instead of trapping it
21 behind an ultrafiltration membrane. The excess reagents are washed away from the
22 absorbed DNA by rinsing the absorptive surface with a wash solution. The solvents
23 used in the wash step must be chosen to avoid loss of the DNA during the wash steps.

1 The basic concept of using a membrane to permit flow of the excess reagent
2 away from the DNA in the wash step can be further combined with the concept of
3 adsorbing the DNA to a surface by incorporating a membrane onto a microfluidic
4 chip. Solvent addition, or washes, may be carried out by the use of electro-osmotic
5 flow. In this particular embodiment, all of the reactions and sample pooling occurs on
6 the chip, permitting high throughput at a lower cost compared to the well plate
7 approach. Within further embodiments, the steps of removing, cleaving, and
8 detecting may be performed in a continuous manner (e.g., as a continuous flow), for
9 example, on a single device which may be automated.

10

11 *Cleavable Tags and Detection*

12 A "tag," according to the present invention, is a chemical moiety that is used
13 to uniquely identify a nucleic acid molecule. In certain preferred embodiments, the
14 nucleic acid molecule is a nucleotide base. In other preferred embodiments, the
15 nucleic acid molecule is a nucleic acid fragment, such as a DNA or an RNA. "Tag"
16 more specifically refers to the tag variable component as well as whatever may be
17 bonded most closely to it.

18 The tags of the present invention further possess one or more of certain
19 characteristic attributes. The tag is preferably distinguishable from all other tags,
20 particularly from other tags used in a particular reaction. The discrimination from
21 other chemical moieties can be based on the chromatographic behavior of the tag
22 (particularly after the cleavage reaction), its spectroscopic or potentiometric
23 properties, or some combination thereof. In addition, the tag is capable of being
24 detected when present at

1 10^{-22} to 10^{-6} mole. The tag is further attachable to the nucleic acid molecule, e.g.,
2 nucleotide base or oligonucleotide, through a "chemical handle" (see U.S. Patent No.
3 6,027,890, incorporated herein by reference) which may attach the tag to the nucleic
4 acid molecule either directly, or through a linker group. In certain preferred
5 embodiments, the tags block primer extension. The tags are further stable toward all
6 manipulations to which they are subjected, including attachment to the nucleic acid
7 molecule and cleavage from the nucleic acid molecule, and any manipulations of the
8 nucleic acid molecule while the tag is attached to it; nor does the tag significantly
9 interfere with the manipulations performed (e.g., hybridization or enzymatic
10 reactions) on the nucleic acid molecule while the tag is attached to it.

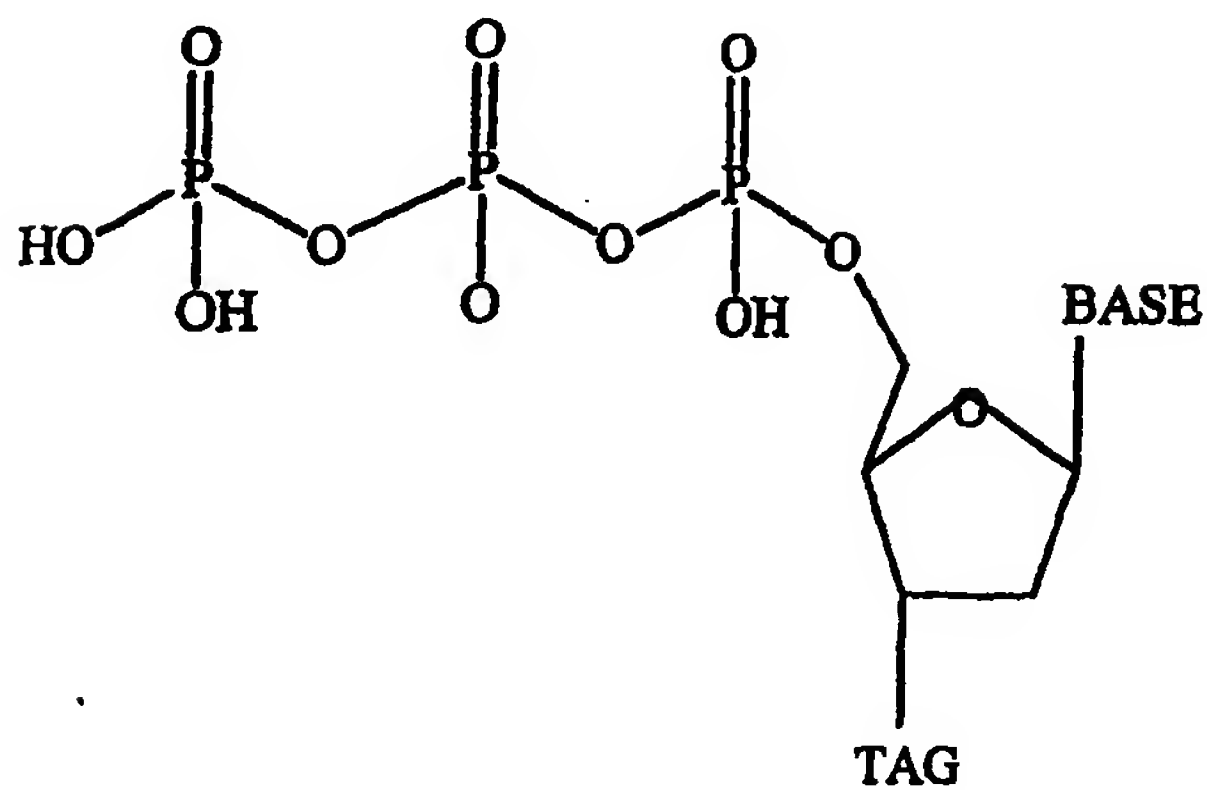
11 The tags of the present invention include any tag that is cleavable by chemical
12 means or by light, and such tags are discussed in detail below. Chemically cleavable
13 tags include tags that are cleavable by an acid or a base. Photo-cleavable tags include
14 tags that are cleavable by a wavelength of light. Other methods of cleavage include
15 oxidation, reduction, enzymatic, electrochemical, heat, and the like.

16 As mentioned above, the tag is further capable of terminating a primer
17 extension reaction. In certain preferred embodiments, the terminating nature of the
18 tag may be due to the nature of the tag itself, for example the structure of the tag, e.g.,
19 a tag that is sufficiently bulky in its structure so that that it prevents addition of any
20 additional bases to the extension product. Alternatively or additionally, the
21 terminating nature of the tag may be due to the placement of the tag on the base.
22 Preferably, the tag is attached to the base so that when the base is added to the
23 growing 3' end of the extension product the tag effectively blocks the extension of the
24 3' end by additional bases, once a tagged base has been added. One such example of

1 a tagged base, wherein the tag is attached directly to the base, that would block

2 extension is shown below.

3

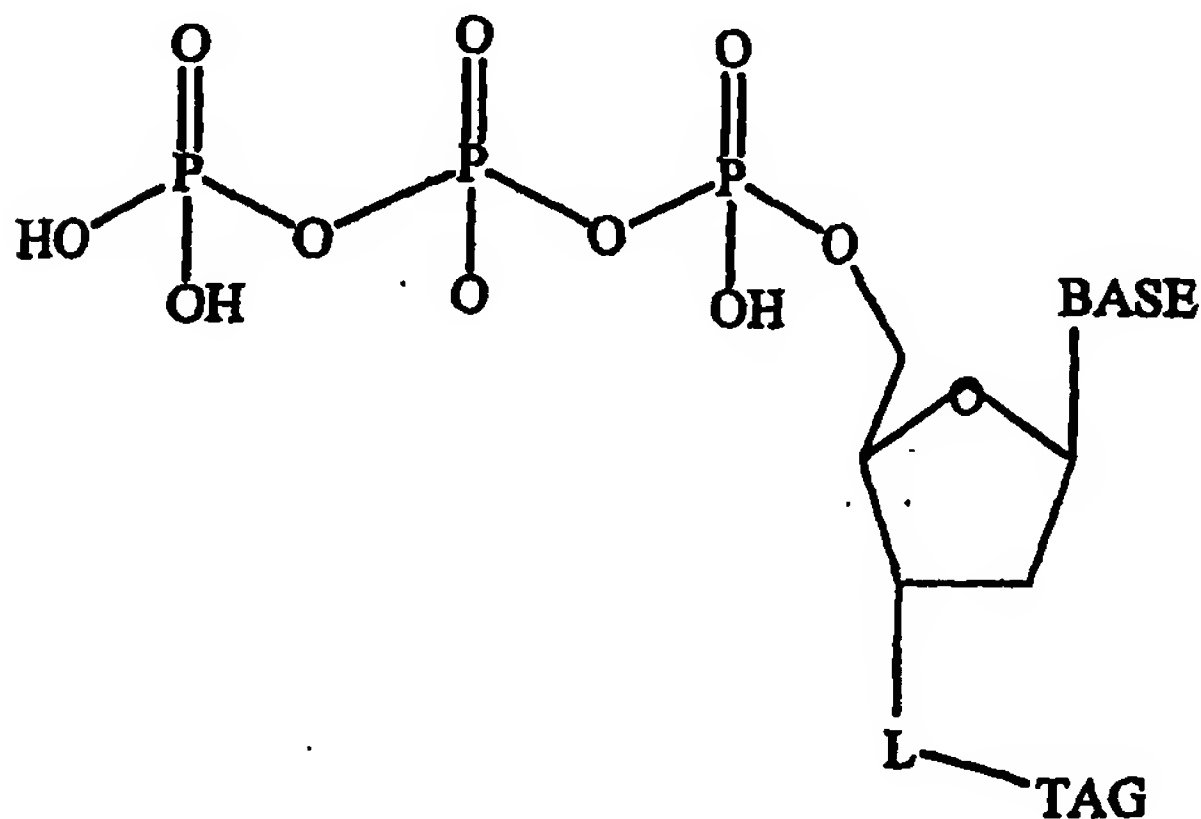


4

5

6 Alternatively, the tag is linked via a labile bond (or labile bonds) to the 3' position of
7 the dNTP, as shown below,

8



9

10

11 wherein:

12 L is the linker.

1 According to the invention, the tag, including the linker in cases where a linker is
2 employed, or other 3' blocking group, are removed to expose the 3' hydroxyl group of
3 the base. Exemplary tags and linkers are described in detail in U.S. Patent 6,027,890,
4 incorporated herein by reference.

5 In light of the availability of numerous tags, any number of tags may be
6 utilized in a given reaction simultaneously, or within different reactions in an array.

7 In certain preferred embodiments, particularly with respect to detection of ligation
8 products, as described below, 4, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 60, 70, 80, 90,
9 100, 150, 200, 250, 300, 350, 400, 450, or greater than 500 different and unique
10 tagged molecules may be utilized within a given reaction simultaneously, wherein
11 each tag is unique for a selected base, oligonucleotide, or other nucleic acid fragment.

12 The characteristics of a variety of well known tags that are amenable to
13 attachment to the bases and nucleic acid molecules of the invention are described in
14 U.S. Patent 6,027,890, incorporated herein by reference. Such tags are detectable,
15 once cleaved from the extended base, by fluorometry, mass spectrometry (MS),
16 infrared (IR) spectrometry, ultraviolet (UV) spectrometry, or potentiostatic
17 amperometry (e.g., utilizing coulometric or amperometric detectors). Mass
18 spectrometry is particularly amendable to multiplexing with mass detection.

19 Representative examples of suitable mass spectrometric techniques include time-of-
20 flight mass spectrometry, quadrupole mass spectrometry, magnetic sector mass
21 spectrometry, and electric sector mass spectrometry. Specific embodiments of such
22 techniques include ion-trap mass spectrometry, electrospray ionization mass
23 spectrometry, ion-spray mass spectrometry, liquid ionization mass spectrometry,
24 atmospheric pressure ionization mass spectrometry, electron ionization mass

1 spectrometry, fast atom bombardment mass spectrometry, MALDI mass
2 spectrometry, photoionization time-of-flight mass spectrometry, laser droplet mass
3 spectrometry, MALDI-TOF mass spectrometry, APCI mass spectrometry, nano-spray
4 mass spectrometry, nebulised spray ionization mass spectrometry, chemical ionization
5 mass spectrometry, resonance ionization mass spectrometry, secondary ionization
6 mass spectrometry, and thermospray mass spectrometry.

7 The following is a list of representative vendors for separation and detection
8 technologies that may be used in the present invention. Perkin Elmer/Applied
9 Biosystems Division (ABI, Foster City, CA) manufactures semi-automated
10 sequencers based on fluorescent-dyes (ABI373) and (ABI377). Analytical Spectral
11 Devices (Boulder, CO) manufactures UV spectrometers. Hitachi Instruments (Tokyo,
12 Japan) manufactures Atomic Absorption spectrometers, Fluorescence spectrometers,
13 LC and GC Mass Spectrometers, NMR spectrometers, and UV-VIS Spectrometers.
14 Perseptive Biosystems (Framingham, MA) produces Mass Spectrometers (VoyagerTM
15 Elite). Bruker Instruments Inc. (Manning Park, MA) manufactures FTIR
16 Spectrometers (Vector 22), FT-Raman Spectrometers, Time of Flight Mass
17 Spectrometers (Reflex IITM), Ion Trap Mass Spectrometer (EsquireTM) and a MALDI
18 Mass Spectrometer. Analytical Technology Inc. (ATI, Boston, MA) makes UV
19 detectors and Diode Array Detectors. Teledyne Electronic Technologies (Mountain
20 View, CA) manufactures an Ion Trap Mass Spectrometer 3DQ DiscoveryTM and the
21 3dQ ApogeeTM). Perkin Elmer/Applied Biosystems Division, (Foster City, CA)
22 manufactures a Sciex Mass Spectrometer (triple quadrupole LC/MS/MS, the API
23 100/300), which is compatible with electrospray. Hewlett-Packard (Santa Clara, CA)
24 produces Mass Selective Detectors (HP 5972A), MALDI-TOF Mass Spectrometers

1 (HP G2025A), Diode Array Detectors, CE units, HPLC units (HP1090), as well as
2 UV Spectrometers. Finnigan Corporation (San Jose, CA) manufactures mass
3 Spectrometers (magnetic sector and four other related mass spectrometers). Rainin
4 (Emeryville, CA) manufactures HPLC instruments.

5 Those skilled in the art will recognize how to apply such devices to the
6 methods of the present invention. Those skilled in the art will further appreciate that
7 devices used to detect pyrosequencing reactions may be adapted to detect and identify
8 the cleaved tags of the invention. For example, the reaction monitoring system
9 described in WO 99/66131, the microfluidic device described in WO 00/40750, the
10 liquid dispensing apparatus described in WO 00/56455, the solid support apparatus of
11 U.S. Patent No. 5,302,509, each of which is incorporated herein by reference, may be
12 adopted for use with the method of the present invention.

13

14 *Automation and High-throughput Sequencing*

15 The DNA sequencing methods of the present invention are fully automatable.
16 Those skilled in the art will recognize that the use of a robot apparatus, where a large
17 number of samples may be rapidly analyzed, may be used for rapid detection and
18 quantification of the tag molecules. Tags to be detected spectrophotometrically may
19 be detected, e.g., by mass spectrometry or fluorescence spectrometry. The use of
20 luminometers, mass spectrometers, and other spectrophotometric devices are well
21 known in the art and described in the literature. The DNA sequencing method of the
22 present invention thus provides an automated approach for high-throughput, non-
23 electrophoretic sequencing procedures that allows for continuous measurement of the
24 progress of the polymerization reaction in real time.

1 In related embodiments, it will be appreciated that multiple samples may be
2 handled in parallel and such parallel handling provides another advantage to the
3 inventive method. In order to obtain high throughput sequence readout, multiple
4 DNA sequencing reactions can be processed in parallel. According to this particular
5 embodiment, the DNA sequencing method of the present invention can be carried out
6 in any of a variety of array formats.

7 For example, a single sequencing reaction of the invention, carried out in a
8 single well and analyzed using flow injection analysis (FIA) has a rate of about one
9 base every six seconds (equivalent to about ten bases per minute and about 600 bases
10 per hour). In order to increase this rate, the DNA sequencing reactions may be
11 multiplexed. For example, multiplexing 25 sequences increases the rate of
12 sequencing to about 15000 bases per hour. Those skilled in the art will recognize the
13 power of multiplexing as it is applicable to any means of detection described herein.
14 The number of DNA samples that can be multiplexed for parallel analysis can range
15 10 to 100, in some cases 100-500, and in yet some other cases, 100-1000 or more
16 DNA samples.

17 In certain preferred embodiments, an array format is used for analysis wherein
18 the DNA samples are distributed over a surface, for example, a microfabricated chip,
19 thereby immobilizing an ordered set of samples in a 2-dimensional format. This
20 allows the analysis of many samples in parallel. According to this embodiment of the
21 invention, the DNA samples are arrayed onto any of a variety available microchips
22 prior to commencing the sequencing reaction. Methods of producing and analyzing
23 DNA arrays are well known in the art and are provided in U.S. Patent No. 6,027,789,
24 incorporated herein by reference.

1 For example, applying the method of the invention to the array format, after
2 primer extension, the tags may be cleaved from the DNA samples on the chip and
3 pooled for analysis using spectrometric or potentiometric techniques (e.g., MALDI-
4 MS). In one particular embodiment of the present invention, an array interrogation
5 system is provided that includes a DNA array generating device, a washing device, a
6 tag cleaving device, a detecting device, and a data processor and analyzer that
7 analyzes data from the detecting device to correlate a tag with a nucleic acid fragment
8 from a sample, as described in U.S. Patent No. 6,027,789, incorporated herein by
9 reference. The arrayed DNA chip has on its surface selected DNA samples of nucleic
10 acid fragments and cleavable tags, e.g., cleavable mass spectrometer tags, attached to
11 the nucleic acid fragments. The arrayed DNA chip is passed through or past a
12 photolytic cleavage device that cleaves the tags from the nucleic acid fragments while
13 still on the DNA chip.

14 After the tags are cleaved, the DNA chip is positioned in an automated micro-
15 array sampling laser device, such as a Matrix Assisted Laser Desorption Ionization
16 (MALDI) instrument. The MALDI instrument is adapted to irradiate and cause
17 desorption of the tags, which are transferred to a detection device, such as a mass
18 spectrometer, wherein tags are identified based upon the difference in molecular
19 weight.

20 Data from the detection device is provided to the data processor and analyzer,
21 which includes a software program that maps the signature of a given tag to a specific
22 sample. The software is able to display the DNA sequence determined and load the
23 sequence information into respective data bases.

1 In an alternative embodiment, the MALDI instrument includes an additional
2 light source that is capable of irradiating the entire DNA chip at a wavelength in the
3 range of 250-360 nm with adjustable intensity, so as to cause the photolytic cleaving
4 of the tags. Accordingly, the cleaving device is incorporated as a component of the
5 MALDI instrument. After cleaving the tags, the MALDI instrument volatized the
6 tags, which are transferred to the detecting device as discussed above.

7 In yet another embodiment, the DNA chip is moved from the DNA array
8 generating device directly to the MALDI instrument. The MALDI instrument
9 includes a laser that emits at a wavelength in the range of approximately 250 to 360
10 nm, inclusive. The laser causes the simultaneous photolytic cleavage of the tag from
11 the nucleic acid fragment along with simultaneous desorption of the tag. The tags are
12 then transferred to the mass spectrometer or other detection device, as discussed
13 above. Accordingly, this alternate embodiment provides photocleavage by the
14 MALDI instrument, so that a separate cleavage device is not needed.

15 If fluorescence sensing is employed in the present invention for detection of
16 the tag, this increases the rate of the sequencing to one base every fifteen seconds
17 (equivalent to about four bases per minute). If 100 sequencing reactions are arranged
18 onto 100 lanes of the chip this yields a rate of about 24000 bases per hour. Similar
19 sequencing rates are achievable with varying cleavage means.

20 Fluorescent tags can be identified and quantitated most directly by their
21 absorption and fluorescence emission wavelengths and intensities. While a
22 conventional spectrofluorometer is extremely flexible, providing continuous ranges of
23 excitation and emission wavelengths (I_{EX} , I_{S1} , I_{S2}), more specialized instruments, such
24 as flow cytometers and laser-scanning microscopes require probes that are excitable at

1 a single fixed wavelength. In contemporary instruments, this is usually the 488-nm
2 line or the argon laser.

3 Radioactive tags may also be applicable to the present invention. Radioactive
4 tags may be detected by, e.g., a CCD detector.

5 In using fluorescent and radioactive tags, the number of different reactions that
6 are simultaneously detectable may be more limited than, e.g., mass tags. For
7 example, the use of four fluorescent molecules, such as commonly employed in DNA
8 sequence analysis, limits analysis to four samples at a time.

9 In certain preferred embodiments, the sample reactions may be pooled on at
10 least one array and the products detected simultaneously. By using a cleavable tag,
11 such as the ones described herein, having a different molecular weight or other
12 physical attribute in each reaction, the entire set of reaction products can be harvested
13 together and analyzed.

14

15 *Applications*

16 The invention in the above embodiments provides a simple and rapid method
17 for sequencing a DNA sample. The methods of the invention both avoid the
18 requirement of separation of the extension product and allows rapid, real-time
19 analysis of the extension reaction. These methods have many applications, which will
20 readily be appreciated by the skilled artisan.

21 To name but a few, the present invention is applicable in the field of forensics
22 (e.g., the identification of individuals and the level of DNA sequence variations);
23 tumor diagnosis (e.g., for detection of viral or cellular oncogenes in a biological
24 sample from a patient); transplantation analyses (e.g., the identification of antigen

1 specific variable DNA sequences from a biological sample); diagnosis of autoimmune
2 diseases, such as juvenile diabetes, arteriosclerosis, multiple sclerosis, rheumatoid
3 arthritis, and encephalomyelitis; genome diagnostics (e.g., the identification of genetic
4 defects or hereditary and acquired genetic diseases in newborns and adults, for
5 example, schizophrenia, manic depression, epilepsy, sickle-cell anemia, thalassemias,
6 α1-antitrypsin deficiency, Lesch-Nyhan syndrome, cystic fibrosis, Duchenn/Becker
7 muscular deficiency, Alzheimer's disease, X-chromosome-dependent mental
8 deficiency, and Huntington's chorea); infectious disease (e.g. detection of viral or
9 microbial infection of a biological sample); mutation detection (e.g., detection of a
10 mutated base in a DNA sample from a biological or artificial source); detection of
11 single nucleotide changes (e.g., a primer hybridizes to a sequence adjacent to a known
12 single nucleotide polymorphism and a dNTP added to the adjacent position is
13 detected and identified).

14 As mentioned above, the method of the present invention may be adapted for
15 use with a ligase instead of a polymerase. One adaptation of this technique is to the
16 oligonucleotide ligation assay, which is used to identify known sequences in very
17 large and complex genomes. To elaborate briefly on the ligase extension reactions
18 described above, the basis of this assay is the ability of a ligase to covalently join two
19 diagnostic oligonucleotides as they hybridize adjacent to one another on a given DNA
20 target. If the sequences at the probe junctions are not perfectly base-paired, the
21 probes will not be joined by the ligase. When tags are used, they are attached to the
22 oligonucleotide, which is ligated to the DNA sample. After a ligation is complete, the
23 tag is cleaved and detected by any of the means described herein (e.g., mass

1 specrometry, infrared spectrophotometry, potentiostatic amperometry, or UV/visible
2 spectrophotometry).

3 In certain preferred embodiment, the DNA sample is amplified prior to
4 exposure to the oligonucleotide ligation assay.

5

6 *Kits*

7 The present invention further provides kits for use in methods of the invention
8 that contain at least the following reagents: a) an oligonucleotide primer suitable for
9 primer extension of a particular DNA template; b) four cdNTPs of adenine, guanine,
10 thymine, and cytosine bases; c) a polymerase; d) a separation means to separate
11 unincorporated dNTPS from the extended DNA template; and e) a cleavage means.

12 In certain embodiments of the invention, a detection means will be provided.

13 However, the detection means may often be provided by the purchaser.

14 In alternative embodiments, if the kit is used for a ligation sequencing reaction
15 assay it may contain at least a) an oligonucleotide primer suitable for primer extension
16 of a particular DNA template; b) at least one tagged oligonucleotide; c) a ligase; d) a
17 separation means to separate unincorporated oligonucleotides from the extended DNA
18 template; and e) a cleavage means. The kit may further provide a detection means.

19 However, the detection means may also be provided by the purchaser.

20 Other embodiments of the invention will be apparent to those skilled in the art
21 from a consideration of the specification or practice of the invention disclosed herein.
22 It is intended that the specification be considered as exemplary only, with the true
23 scope and spirit of the invention being indicated by the following claims.

1 What is claimed is:

2 1. A method of determining the sequence of a nucleic acid, comprising steps of:

3 (a) hybridizing an oligonucleotide to a single stranded DNA, wherein
4 the oligonucleotide is complementary to at least a portion of the single
5 stranded DNA;

6 (b) providing a DNA polymerase and four deoxynucleotide
7 triphosphates (dNTPs) comprising dATP, dGTP, dCTP, and dTTP,
8 wherein each dNTP is 3'-end labeled with a cleavable tag (cdNTP) that
9 distinguishes it from the other cdNTPs;

10 (c) extending the single stranded DNA hybridized to the
11 oligonucleotide by one complementary end-labeled cdNTP in a
12 polymerase extension reaction, wherein the tag on the extended cdNTP
13 blocks further extension by the DNA polymerase;

14 (d) cleaving the tag from the complementary cdNTP; and

15 (e) detecting the tag, thereby identifying the complementary dNTP.

16 2. The method of claim 1, further comprising the step of removing excess
17 cdNTPs that are not extended onto the single stranded DNA.

18 3. The method of claim 1, further comprising the step of repeating steps (a)
19 through (e) on the sample of single stranded DNA.

1 4. The method of claim 1, wherein the cleavable tags are cleavable by chemical
2 cleavage.

3 5. The method of claim 4, wherein the cleavable tags are acid cleavable tags.

4 6. The method of claim 4, wherein the cleavable tags are base cleavable.

5 7. The method of claim 1, wherein the tags are photocleavable.

6 8. The method of claim 1, wherein the tag is a fluorescent tag.

7 9. The method of claim 1, wherein the tag is a mass tag.

8 10. A method of determining the sequence of a nucleic acid, comprising steps of:

9 (a) hybridizing a complementary oligonucleotide to a single stranded DNA,
10 wherein the oligonucleotide is 3'-end labeled with one or more cleavable tags
11 that distinguishes it from other oligonucleotides;

12 (b) cleaving the one or more tags from the ligated complementary
13 oligonucleotide; and

14 (c) detecting the one or more tags.

15 11. The method of claim 10, wherein the hybridizing of the complementary
16 oligonucleotide occurs adjacent to a primer.

17 12. The method of claim 11, further comprising ligating the hybridized
18 oligonucleotide to the primer.

- 1 13. The method of claim 12, further comprising the step of removing excess
- 2 oligonucleotides that are not ligated onto the single stranded DNA.
- 3 14. The method of claim 12, further comprising the step of repeating steps (a)
- 4 through (c) on the single stranded DNA.
- 5 15. The method of claim 10, wherein one or more tags are cleaved by chemical
- 6 cleavage.
- 7 16. The method of claim 15, wherein the cleavable tags are acid cleavable tags.
- 8 17. The method of claim 15, wherein the cleavable tags are base cleavable.
- 9 18. The method of claim 10, wherein the tags are photocleavable.
- 10 19. The method of claim 10, wherein the tag is a fluorescent tag.
- 11 20. The method of claim 10, wherein the tag is a mass tag.

12

INTERNATIONAL SEARCH REPORT

International application No. PCT/US02/14198

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) :C12Q 1/68, C12P 19/34, C07H 21/04

US CL :Please See Extra Sheet.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 91.2; 586/24.38, 25.8, 27; 985/77,78

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
None

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

stn searched, uspatfull, medline, biosis, caplus

key words: DNA sequencing, ligation, cleavable, tag, labeled, mass, fluorescent, polymerase, solid support

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 5,302,509 A (CHEESEMAN) 12 April 1994, col. 2, lines 15-49, col. 6, lines 1-12.	1-9
Y		---
X	US 5,552,278 (BRENNER), 03 September 1996, col. 6, lines 45-59 and 64-67 and col. 7, lines 11-14.	10-14 and 19-20

Further documents are listed in the continuation of Box C. See patent family annex.

* Special categories of cited documents:	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"A" document defining the general state of the art which is not considered to be of particular relevance	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"B" earlier document published on or after the international filing date	"Y"	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"&"	document member of the same patent family
"O" document referring to an oral disclosure, use, exhibition or other means		
"P" document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search

15 JUNE 2002

Date of mailing of the international search report

03 JUL 2002

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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/14193

A. CLASSIFICATION OF SUBJECT MATTER:
US CL :

435/6, 91.2; 536/24.88, 25.3, 97; 935/77,78